

On page 4, please replace the second through fourth paragraphs with the following rewritten paragraphs:

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--Figures 1A-1D show the cDNA sequence and corresponding deduced amino acid sequence of the mature NTT polypeptide. The standard one-letter abbreviations are utilized to represent the amino acid residues in the polypeptide sequence shown in Figures 1A-1D.

In accordance with an aspect of the present invention, there is provided an isolated nucleic acid (polynucleotide) which encodes for the mature polypeptide having the deduced amino acid sequence of Figures 1A-1D, collectively, or for the mature polypeptide encoded by the cDNA of the clone deposited as ATCC Deposit No. 75713 on March 18, 1994. This deposit is a biological deposit with the American Type Culture Collection (ATCC), 10801 University Blvd., Manassas, Virginia 20110-2209, USA.

The polynucleotide of this invention was discovered in a cDNA library derived from a human fetal brain. It is structurally related to the neurotransmitter transporter family. It contains an open reading frame encoding a protein of about 727 amino acid residues. The protein exhibits the highest degree of homology to a rat neurotransmitter transporter (NTT4) with 94% identity and 96% similarity over the entire amino acid sequence.

The polynucleotide of the present invention may be in the form of RNA or in the form of DNA, which DNA includes cDNA, genomic DNA, and synthetic DNA. The DNA may be double-stranded or single-stranded, and if single stranded may be the coding strand or non-coding (anti-sense) strand. The coding sequence which encodes the mature polypeptide may be identical to the coding sequence shown in Figures 1A-1D, collectively, or that of the deposited clone or may be a different coding sequence which coding sequence, as a result of the redundancy or degeneracy of the genetic code, encodes the same, mature polypeptide as the DNA of Figures 1A-1D, collectively, or the deposited cDNA.--

**[Starting on page 4, please replace the paragraph bridging pages 4 and 5 with the following rewritten paragraph:]**

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--The polynucleotide which encodes for the mature polypeptide of Figures 1A-1D, collectively, or for the mature polypeptide encoded by the deposited cDNA may include: only the coding sequence for the mature polypeptide; the coding sequence for the mature polypeptide and additional coding sequence such as a leader or secretory sequence or a proprotein sequence; the coding sequence for the mature polypeptide (and optionally additional coding sequence) and non-coding sequence, such as introns or non-coding sequence 5' and/or 3' of the coding sequence for the mature polypeptide.--

**[On page 5, please replace the second through fourth full paragraph with the following rewritten paragraphs:]**

--The present invention further relates to variants of the hereinabove described polynucleotides which encode for fragments, analogs and derivatives of the polypeptide having the deduced amino acid sequence of Figures 1A-1D, collectively, or the polypeptide encoded by the cDNA of the deposited clone. The variant of the polynucleotide may be a naturally occurring allelic variant of the polynucleotide or a non-naturally occurring variant of the polynucleotide.

Thus, the present invention includes polynucleotides encoding the same mature polypeptide as shown in Figures 1A-1D, collectively, or the same mature polypeptide encoded by the cDNA of the deposited clone as well as variants of such polynucleotides which variants encode for a fragment, derivative or analog of the polypeptide of Figures 1A-1D, collectively, or the polypeptide encoded by the cDNA of the deposited clone. Such nucleotide variants include deletion variants, substitution variants and addition or insertion variants.

As hereinabove indicated, the polynucleotide may have a coding sequence which is a naturally occurring allelic variant of the coding sequence shown in Figures 1A-1D, collectively, or of the coding sequence of the deposited clone. As known in the art, an allelic variant is an alternate form of a polynucleotide sequence which may have a substitution, deletion or addition of one or more nucleotides, which does not substantially alter the function of the encoded polypeptide.--

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01013	01012	01011	01010	01009	01008	01007	01006	01005	01004	01003	01002	01001	01000	00999	00998	00997	00996	00995	00994	00993	00992	00991	00990	00989	00988	00987	00986	00985	00984	00983	00982	00981	00980	00979	00978	00977	00976	00975	00974	00973	00972	00971	00970	00969	00968	00967	00966	00965	00964	00963	00962	00961	00960	00959	00958	00957	00956	00955	00954	00953	00952	00951	00950	00949	00948	00947	00946	00945	00944	00943	00942	00941	00940	00939	00938	00937	00936	00935	00934	00933	00932	00931	00930	00929	00928	00927	00926	00925	00924	00923	00922	00921	00920	00919	00918	00917	00916	00915	00914	00913	00912	00911	00910	00909	00908	00907	00906	00905	00904	00903	00902	00901	00900	00899	00898	00897	00896	00895	00894	00893	00892	00891	00890	00889	00888	00887	00886	00885	00884	00883	00882	00881	00880	00879	00878	00877	00876	00875	00874	00873	00872	00871	00870	00869	00868	00867	00866	00865	00864	00863	00862	00861	00860	00859	00858	00857	00856	00855	00854	00853	00852	00851	00850	00849	00848	00847	00846	00845	00844	00843	00842	00841	00840	00839	00838	00837	00836	00835	00834	00833	00832	00831	00830	00829	00828	00827	00826	00825	00824	00823	00822	00821	00820	00819	00818	00817	00816	00815	00814	00813	00812	00811	00810	00809	00808	00807	00806	00805	00804	00803	00802	00801	00800	00799	00798	00797	00796	00795	00794	00793	00792	00791	00790	00789	00788	00787	00786	00785	00784	00783	00782	00781	00780	00779	00778	00777	00776	00775	00774	00773	00772	00771	00770	00769	00768	00767	00766	00765	00764	00763	00762	00761	00760	00759	00758	00757	00756	00755	00754	00753	00752	00751	00750	00749	00748	00747	00746	00745	00744	00743	00742	00741	00740	00739	00738	00737	00736	00735	00734	00733	00732	00731	00730	00729	00728	00727	00726	00725	00724	00723	00722	00721	00720	00719	00718	00717	00716	00715	00714	00713	00712	00711	00710	00709	00708	00707	00706	00705	00704	00703	00702	00701	00700	00699	00698	00697	00696	00695	00694	00693	00692	00691	00690	00689	00688	00687	00686	00685	00684	00683	00682	00681	00680	00679	00678	00677	00676	00675	00674	00673	00672	00671	00670	00669	00668	00667	00666	00665	00664	00663	00662	00661	00660	00659	00658	00657	00656	00655	00654	00653	00652	00651	00650	00649	00648	00647	00646	00645	00644	00643	0064
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The terms "fragment," "derivative" and "analog" when referring to the polypeptide of Figures 1A-1D, collectively, or that encoded by the deposited cDNA, means a polypeptide which retains essentially the same biological function or activity as such polypeptide. Thus, an analog includes a proprotein which can be activated by cleavage of the proprotein portion to produce an active mature polypeptide.--

Starting on page 7, please replace the paragraph bridging pages 7 and 8 with the following rewritten paragraph:

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--The fragment, derivative or analog of the polypeptide of Figures 1A-1D, collectively, or that encoded by the deposited cDNA may be (i) one in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue) and such substituted amino acid residue may or may not be one encoded by the genetic code, or (ii) one in which one or more of the amino acid residues includes a substituent group, or (iii) one in which the mature polypeptide is fused with another compound, such as a compound to increase the half-life of the polypeptide (for example, polyethylene glycol), or (iv) one in which the additional amino acids are fused to the mature polypeptide, such as a leader or secretory sequence or a sequence which is employed for purification of the mature polypeptide or a proprotein sequence. Such fragments, derivatives and analogs are deemed to be within the scope of those skilled in the art from the teachings herein.--

Starting on page 23, please replace the paragraph bridging pages 23-25 with the following rewritten paragraph:

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The DNA sequence encoding for NTT, ATCC # 75713 is initially amplified using PCR oligonucleotide primers corresponding to the 5' and sequences of the processed NTT protein (minus the signal peptide sequence) and the vector sequences 3' to the NTT gene. Additional nucleotides corresponding to NTT were added to the 5' and 3' sequences respectively. The 5' oligonucleotide primer has the sequence GACTAAAGCTTGGCATCAATGCCGAAGAAC (SEQ ID NO:3) contains a Hind III restriction enzyme site followed by 18 nucleotides of NTT coding sequence. The 3' sequence GAACTTCTAGAGCAGTGGTCACAGCTCAG (SEQ ID NO:4) contains complementary sequences to Xba I site and is followed by 18 nucleotides of NTT sequence. The restriction enzyme sites correspond to the restriction enzyme sites on the bacterial expression vector pQE-9. (Qiagen, Inc. 9259 Eton Avenue, Chatsworth, CA, 91311). pQE-9 encodes antibiotic resistance (Amp<sup>r</sup>), a bacterial origin of replication (ori), an IPTG-regulatable promoter operator (P/O), a ribosome binding site (RBS), a 6-His tag and restriction enzyme sites. pQE-9 was then digested with Hind III and Xba I. The amplified sequences were ligated into pQE-9 and were inserted in frame with the sequence encoding for the histidine tag and the RBS. The ligation mixture was then used to transform *E. coli* strain M15/rep 4 available from Qiagen under the trademark M15/rep 4 by the procedure described in Sambrook, J. et al., Molecular Cloning: A Laboratory Manual, Cold Spring Laboratory Press, (1989). M15/rep4 contains multiple copies of the plasmid pREP4, which expresses the lacI repressor and also confers kanamycin resistance (Kan<sup>r</sup>). Transformants are identified by their ability to grow on LB plates and ampicillin/kanamycin resistant colonies were selected. Plasmid DNA was isolated and confirmed by restriction analysis. Clones containing the desired constructs were grown overnight (O/N) in liquid culture in LB media supplemented with both Amp (100 ug/ml) and Kan (25 ug/ml). The O/N culture is used to inoculate a large culture at a ratio of 1:100 to 1:250. The cells were grown to an optical density 600 (O.D.<sub>600</sub>) of between 0.4 and 0.6. IPTG ("Isopropyl-B-D-thiogalacto pyranoside") was then added to a final concentration of 1 mM. IPTG induces by inactivating the lacI repressor, clearing the P/O leading to increased gene expression. Cells were grown an extra 3 to 4 hours. Cells were then harvested by

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Starting on page 25, please replace the paragraph bridging pages 25 and 26 with the following rewritten paragraph:

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--The DNA sequence encoding for NTT, ATCC # 75713, was constructed by PCR on the original EST cloned using two primers: the 5' primer GACTAAGATCTGCCACCATGCCGAAGAACAGCAAAGTG (SEQ ID NO:5) contains a Bgl II site followed by 21 nucleotides of NTT coding sequence starting from the initiation codon; the 3' sequence GAACTGATATCGCAGTGGTCACAGCTCAG (SEQ ID NO:6) contains complementary sequences to EcoR V site, translation stop codon, and the last 18 nucleotides of the NTT coding sequence. Therefore, the PCR product contains a Bgl II site, NTT coding sequence followed by a translation termination stop codon, and an EcoR V site. The PCR amplified DNA fragment and the vector, pcDNAI/Amp, were digested with Bgl II and EcoR V. The ligation mixture was transformed into E. coli strain SURE (available from Stratagene Cloning Systems, 11099 North Torrey Pines Road, La Jolla, CA 92037) the transformed culture was plated on ampicillin media plates and resistant colonies were selected. Plasmid DNA was isolated from transformants and examined by restriction analysis for the presence of the correct fragment. For expression of the recombinant NTT, COS cells were transfected with the expression vector by DEAE-DEXTRAN method. (J. Sambrook, E. Fritsch, T. Maniatis, Molecular Cloning: A Laboratory Manual, Cold Spring Laboratory Press, (1989)). The expression of the NTT HA protein was detected by radiolabelling and immunoprecipitation method. (E. Harlow, D. Lane, Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory Press, (1988)). Cells were labelled for 8 hours with <sup>35</sup>S-cysteine two days post transfection. Culture media were then collected and cells were lysed with detergent (RIPA buffer (150 mM NaCl, 1% NP-40, 0.1% SDS, 1% NP-40, 0.5% DOC, 50mM Tris, pH 7.5). (Wilson, I. et al., Id. 37:767 (1984)). Both cell lysate and culture media were precipitated with a HA specific monoclonal antibody. Proteins precipitated were analyzed on 15% SDS-PAGE gels.--

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1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54	55	56	57	58	59	60	61	62	63	64	65	66	67	68	69	70	71	72	73	74	75	76	77	78	79	80	81	82	83	84	85	86	87	88	89	90	91	92	93	94	95	96	97	98	99	100

**In the Drawings:**

PF116D1C1